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Ultrahigh-sensitivity label-free optical fiber biosensor based on a tapered singlemode- no core-singlemode coupler for *Staphylococcus aureus* detection

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Abstract

An ultra-high sensitivity label-free optical fiber biosensor for inactivated *Staphylococcus aureus* (*S. aureus*) detection is proposed and investigated in this study, with additional advantages of robust and stability compared to traditional tapered fiber structure. The proposed fiber biosensor is based on a tapered singlemode- no coresinglemode fiber coupler (SNSFC) structure, where the no core fiber was tapered to small diameter (taper-waist diameter of about 10 μ m) and functionalized with the pig immunoglobulin G (IgG) antibody for detection of *S. aureus*. The measured maximum wavelength shift of the sensor for an *S. aureus* concentration of 7×10¹ CFU/ml (colony forming unit per milliliter) is 2.04 nm, which is equivalent to a limit of detection (LOD) of 3.1 CFU/ml (a highest LOD reported so far for optical fiber biosensors), considering the maximum wavelength variation of the sensor in phosphate buffered saline (PBS) is ±0.03 nm over 40 minutes, where 3 times of maximum wavelength variation (3×0.03=0.09 nm) is defined as measurement limit. The response time of the developed fiber sensor is less than 30 minutes. The ultra-sensitive biosensor has potential to be widely applied to various areas such as disease, medical diagnostics and food safety inspection.

Keywords: Optical fiber sensor, biosensor, Staphylococcus aureus (S. aureus), foodborne pathogens.

1. Introduction

Staphylococcus aureus (S. aureus), firstly discovered by Dr. Alexander Ogston in 1880, is a type of spherical bacteria with a diameter of about 0.8 µm and arranged like a cluster of grapes under the microscope without spores [1]. S. aureus is one of the most common foodborne pathogens and often parasitic on human and animal skin, nasal cavity, stomach, air and sewage, which can produce variety of toxins causing diseases, such as pneumonia, pus infections and pericarditis, etc [2-5]. In order to avoid the diseases caused by S. aureus, it is crucial to achieve rapid and highly sensitive detection of S. aureus before the contaminated food is consumed. The traditional detection methods for the detection of S. aureus are culture-based assay, which normally requires several days to culture the bacteria before the results can be obtained. Most foods are contaminated with fewer pathogen cells [normally less than 100 CFU/g (Colony Forming Unit per gram)]. Therefore, an initial enrichment of the sample is needed with the traditional methods. However, the products are minimally processed and have an inherently short shelf life, which limited the application of these traditional methods. Hence, there is an urgent need to develop a high sensitivity, rapid detection method for foodborne pathogens. In recent years, some rapid detection technologies have been developed, such as polymerase chain reaction (PCR) [6-7], enzyme linked immunosorbent assay (ELISA) [8] and nucleic acid-based molecular biology methods [9]. For example, Majumdar et al reported an electrochemical biosensor for detection of S. aureus by depositing antibody on the surface modified platinum electrode, which achieved a limit of detection (LOD) of 10 CFU/ml. However, these electrochemical biosensors have disadvantages of generating false current values due to the

existence of different electro-active compounds, thus often leading to interferences and hence measurement errors [10]. In 2017, Menti *et al* developed a mass-based biosensor based on the magneto-elastic method for *S. aureus* detection. However, this technique has been limited with a relatively poor LOD of 10^4 CFU/ml and complicated operation processes [11-12].

In comparison with these techniques, optical biosensors have unique advantages such as high sensitivity, simple fabrication processes, capability for real time monitoring and robust to environments, which are widely investigated for detection of pathogens [13]. Depending on different working principles, optical biosensors can be divided into different categories, including fiber grating [14-15], surface plasmon resonance (SPR) [16], tapered optical fiber [17-18], surface enhanced Raman spectroscopy (SERS) [19], colorimetric sensor [20], fluorescence [21], U-bent fiber [22] and optical fiber interferometry [23]. For example, Tripathi et al adopted a long period grating modified with T4 bacteriophage to detect Escherichia coli (E.coli) and achieved an LOD of 10³ CFU/ml [24]. A tapered fiber biosensor was developed to specifically identify dead E. coli O157:H7 and obtained an LOD of 10⁴ CFU/ml [25]. Janczuk-Richter et al demonstrated a long period grating based biosensor for T7 bacteriophage detection and achieved an LOD less than 5×10^3 CFU/ml [26]. Abdelhamid and Wu et al utilized fluorescence optical biosensors to detect S. aureus and achieved an LOD of 2×10^2 CFU/ml [27]. Most recently, Kumar et al developed an ultra-highly sensitive biosensor based on tapered no-core fiber structure for human chorionic gonadotropin (hCG) detection with an LOD of 0.0001 mIU/ml [28]. The tapered fiber interferometer provides an ultrahigh sensitivity, however due to a very small diameter (~10 µm) of the taper waist, this type of sensor is fragile with relatively poor stability. In this paper, we propose a novel tapered fiber biosensor based on tapered no-core fiber coupler structure, which has a much larger taper waist diameter (double waist diameter of a single taper fiber and hence robust with a better stability) without sacrificing the sensitivity, and applied it for detection of S. aureus.

2. Sensor structure

Figure1 shows a schematic diagram of the tapered singlemode-no core-singlemode fiber coupler (SNSFC) structure. When the light is transmitted from input singlemode fiber (SMF) to the tapered no core fiber (NCF), it will excite multiple cladding modes and coupled to the other NCF which is physically contacted with the input NCF. The light will then be coupled to the two output SMF fibers. The multiple cladding modes in the NCF section will interact with the surrounding environment, thus resulting in a wavelength shift or power variation for the tapered SNSFC structure. Once the wavelength shift or the power variation is calibrated, the change of surrounding environment will be determined.



Fig. 1. Schematic diagram of SNSFC fiber structure

3. Experiments

3.1 The tapered SNSFC structure for refractive index (RI) measurement.

In the experiments, two singlemode-no core-singlemode (SNS) fiber sensor structures were firstly prepared by fusion splicing a short section of NCF (15 mm length) between two SMFs. The NCF used in the experiments is FG125LA purchased from Thorlabs. The fabricated two SNS fiber structures were then aligned and twisted two turns between two fiber holders with a distance of 80 mm, which enables the two SNS fiber structures having good physical contact. The NCF sections of the two SMS fiber structures were then heated to circa 1200 °C that can soften the NCFs, and the two ends of the SNS were pulled by two computer-controlled translation stages for thinning the NCF to form a tapered SNSFC structure. Two tapered SNSFCs with different taper waist diameters

(58.6 and 10.4 μ m) were prepared for RI sensitivity measurement. Figure 2(a) shows wavelength shifts of the sensors with two different taper waist diameters at the RI range of 1.33. The sensor with a smaller diameter of 10.4 μ m has a higher RI sensitivity of 1523.333 nm/RIU, compared to that of 233.977 nm/RIU for the sensor with a taper waist diameter of 58.6 μ m. Therefore, the fiber sensor with a taper diameter of 10.4 μ m is selected for further experiments for detection of *S. aureus* (provided from State Key Lab Food Science and Technology, Nanchang University, China). Since *S. aureus* normally survived in phosphate buffered saline (PBS; Catalog number: SH30256.01, pH=7.4, purchased from GE Healthcare Life Sciences) environment, the stability of the biosensor in PBS was firstly tested. Figure 2(b) shows the tapered SNSFC sensors have good stability with maximum wavelength variations of ±0.03 nm over 40 minutes in the PBS.



Fig. 2. (a) Wavelength shift of the tapered SNSFC vs. RI with two different taper waist diameters (58.6 and 10.4 μ m) and (b) stability of the tapered SNSFC sensor in PBS with a taper waist diameter of 10.4 μ m.



Fig. 3. A schematic diagram of fiber surface modification process: (i) treat with silane reagent to create carboxyl group; (ii) generate NHS active ester with EDC/NHSS; (iii) immobilize pig immunoglobulin G (IgG) antibodies on the fiber sensor surface; (iv) block unbind sites with BSA; (v) specific bind with *S. aureus*

3.2 Functionalization of the tapered SNSFC sensor

In order to enable the fiber sensor for *S. aureus* detection, the sensor needs to be functionalized with pig IgG antibody (product code: b1108, purchased from BEIJING BERSEE SCIENCE AND TECHNOLOGY CO.LTD, http://www.berseebio.com), which can specifically bind with target *S. aureus* [29-32]. The functionalization process is illustrated below:

i. The fiber sensor is immersed in a solution of 5% silane reagent (3-(3-triethoxysilylpropyl)oxolane-2,5dione) (product code:T195932-5g, purchased from aladdin) in ethanol for 4 hours at room temperature to create a carboxyl group on the fiber sensor surface.

- ii. After cleaning the above fiber with deionized water, the fiber is then immersed into freshly prepared pH buffer(pH=6.0, product code: P118681 and P118682, purchased from aladdin) containing 1-(3-Dimethylaminopropyl)-3-ethylcar-bodiimide hydrochloride (EDC, product code: 39141134, purchased from Sinopharm Chemical Reagent Co., Ltd) for 30 minutes, followed by immediately immersed in a mixed solution of EDC and hydroxy-2,5-dioxopyrolidine-3-sulfonicacid sodium salt (NHSS, product code: H109337, purchased from aladdin) for 30 minutes.
- iii. The above fiber sensor is washed with PB buffer and then immersed into a pig IgG antibody solution (in PBS buffer) for 4 hours. This process enables the pig IgG antibody be immobilized on the surface of fiber sensor.
- iv. After washing the above fiber with a PBS, the above fiber sensor is then immersed into 1% Bovine serum albumin (BSA, product code: A119741, purchased from aladdin) in PBS for 2 hours at room temperature, then the fiber sensor is cleaned with PBS before being used for detection of *S. aureus*.

A schematic diagram of the fiber sensor surface functionalization process is illustrated in Fig. 3(b): i-iv, which corresponds to above four steps, respectively. The functionalized fiber sensor will specifically bind with S. *aureus* as shown in Fig. 3(b): v.

Figures 4(a) and (b) illustrated scanning electron microscope (SEM) images of the functionalized fiber sensor bind with *S. aureus* (the sphere in Fig. 4).





3.3 Detection of *S. aureus*

Figure 5 shows a schematic diagram of the experimental setup for detection of *S. aureus* by using the developed fiber sensor, where a broadband light source (BBS, SC-5-FC) is connected to one of the input SMFs and an optical spectrum analyzer (OSA, YOKOGAWA AQ6370D) is used to measure spectral response of the tapered SNSFC sensor. The fiber coupler sensor section was immersed in a rectangular container (44 $mm \times 5 mm \times 5$ mm), in which different concentrations of *S. aureus* samples were added in sequence (from low to high: 7×10^1 CFU/ml, 7×10^2 CFU/ml, 7×10^3 CFU/ml, 7×10^4 CFU/ml) during the tests. Before changing each concentration of *S. aureus* sample, the functionalized sensor was immersed into PBS buffer for 20 minutes to wash out the nonspecific bind *S. aureus*. The fiber sensor was then immersed into *S. aureus* sample for 30 minutes to allow the pig IgG antibodies specifically capture and bind with *S. aureus*, which effectively introduces change of the surrounding RI and coating thickness of the fiber sensor, resulting in change of spectral responses.



Fig. 5. Schematic diagram of the experimental setup of the tapered SNSFC for S. aureus detection

4. Results and discussion

Investigation of capture layer concentration

The spectral response of the functionalized sensor (sensor surface functionalized with 50 μ g/ml pig IgG antibody) at different durations are shown in Fig. 6(a) by immersing the fiber sensor within *S. aureus* solution with a concentration of 7×10¹ CFU/ml. The dip wavelength is shifted monotonically to short wavelength side as the time is increased up to 30 minutes.

The influence of the concentration of pig IgG antibody on the performance of the sensor was studied using three different tapered SNSFC structures (fabricated under the same conditions) functionalized with three different concentrations (25 µg/ml, 50 µg/ml and 200 µg/ml) of pig IgG antibody to detect *S. aureus* samples with different concentrations of 7×10^1 CFU/ml, 7×10^2 CFU/ml, 7×10^3 CFU/ml, 7×10^4 CFU/ml in sequence, respectively. Figure 6(b) summarized the relationships between time and dip wavelength shifts of the three fiber biosensors immersed in different concentrations of *S. aureus* samples.

It can be clearly seen from Fig. 6(b) that, for all the three fiber biosensors, as the immerse time increases, the dip wavelength shifts monotonically to the short wavelength side, which is due to the binding process of S. aureus onto the functionalized fiber sensor surface. It also shows that the higher the S. aureus concentration, the larger the dip wavelength shift. Whereas the dip wavelength shift is mainly observed in the initial 20 minutes, after which the spectral responses become stabilized, indicating that the binding process is mainly taken place in the first 20 minutes. During the binding process, the shift of wavelength shows an exponential relationship with immersing time, which is consistent with the kinetic response of the immune behaviour reported in the literature [33]. At the same concentration of S. aureus, the sensor functionalized with a higher concentration of pig IgG antibody has a much larger dip wavelength shift and hence a higher sensitivity. The possible reason is that a higher concentration of pig IgG antibody has better capability to capture S. aureus and hence result in a larger wavelength shift for the same concentration of S. aureus. Reproducibility of the biosensors was investigated by fabricating 15 tapered SNSFC fiber structure with same fabrication parameters. The fabricated 15 tapered SNSFC structures were then functionalized with 25 μ g/ml, 50 μ g/ml and 200 μ g/ml pig IgG antibody, where each concentration of pig IgG antibody is used to modify 5 tapered SNSFC structures. The measurement results were shown in Fig. 6(c), which clearly show that the tapered SNSFC biosensor has a relatively good reproducibility. The average dip wavelength shifts of fiber biosensors modified with 25 µg/ml, 50 µg/ml and 200 µg/ml were 0.25, 0.56, 0.87, 1.20; 0.70, 1.76, 3.09, 4.22 and 2.04 nm, 3.73 nm, 5.25 nm, 7.58 nm for different concentrations (7×10¹ CFU/ml, 7×10² CFU/ml, 7×10³ CFU/ml, 7×10⁴ CFU/ml) of S. aureus. It is noted that, in the above calculation of average dip wavelength shift, the accumulated wavelength variations in PBS solution as shown in Fig. 6(b) weren't counted. Figure 6(c) also shows that the fiber biosensor has largest error bar at the lowest concentration of 7×10^1 CFU/ml compared to that of highest concentration of 7×10^4 CFU/ml. This is possibly because that number of S. aureus of one colony isn't certain and the higher concentration of S. aureus sample will have better average effect compared to that of lower concentration of S. aureus sample, and thus smaller measurement difference with different fiber biosensors. When the concentration of S. aureus is 7×10^1 CFU/ml, the sensor functionalized with pig IgG antibody concentration of 200 µg/ml has the largest average wavelength shift of 2.04 nm. Since the wavelength variation of the fiber sensor in PBS is within ± 0.03 nm, the LOD for S. aureus reachs 3.1 CFU/ml provided the measurement limit is defined as 3 times of the wavelength variation (0.09 nm).

The specificity of the biosensor was further investigated by immersing three identical sensors (functionalized with 50 µg/ml pig IgG antibody) into three different analytes (varying from large bacteria *E.Coli* to small protein hCG), namely 4×10^6 CFU/ml *E. coli*, 10 mg/ml BSA and 50 mIU/ml hCG and each analytes were tested five times. Experimental results combined with measurement of *S. aureus* with concentration of 7×10^4 CFU/ml are shown in Fig.6 (e). The average dip wavelength shifts of identical sensors were observed to be 4.22; 0.14; 0.23; and 0.42 nm for the *S. aureus*, *E. coli*, BSA and hCG respectively, indicating that the biosensor has relatively good specificity and reproducibility.



Fig. 6. (a) Spectral responses of the tapered SNSFC sensor functionalized with 50 μ g/ml pig IgG antibody for detection of *S. aureus* with concentration of 7×10¹ CFU/ml; (b) comparison of different capture layer concentrations (25 μ g/ml, 50 μ g/ml, 200 μ g/ml) for *S. aureus* detection; (c) reproducibility and (d) specificity results of the biosensor.

Table 1	A summary	of optical	biosensor	techniques	used for the	detection of	pathogenic	bacteria
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Transducer	Pathogenic bacteria	LOD	Reference	
Long-period fiber gratings (LPFGs)	S. aureus	224 CFU/ml	[34] (2019)	
A fiber optic SPR immunosensor with functionalized molybdenum disulfide (MoS2) nanosheets	E. coli	94 CFU/mL	[35] (2019)	
Fluorescent (optical) bioprobe	S. aureus 85 CFU/mL		[36] (2019)	
Multi-column capillary biosensor based on Fenanocluster amplification and smart phone imaging	Salmonella Typhimurium (S. Typhimurium)	14 CFU/mL	[37] (2019)	
Multimode microfiber	E. coli	10 ³ CFU/ml	[38] (2018)	
Single mode-tapered multimode- single mode (SMS)	S. Typhimurium	247 CFU/mL	[39] (2018)	
Microfluidic biosensor based on fluorescence labelling and smartphone video processing	S. Typhimurium	58 CFU/mL	[40] (2019)	
Fluorescence imaging biosensor by smartphone	S. aureus	10 CFU/mL	[41] (2018)	
Microcavity in-line Mach-Zehnder	Live E. coli	100 CFU/mL	[42] (2018)	

interferometer			
Tapered SNSFC biosensor	S. aureus	3.1 CFU/mL	Proposed method in this paper

The performance comparison between our newly developed fiber biosensor and those of recently published optical biosensor techniques for the detection of bacteria are shown in Table 1. The LOD of our proposed sensor is as low as 3.1 CFU/ml, which is the best result reported so far by using optical fiber sensors, demonstrating the significant advantage of our proposed sensor.

As for the lifetime of the sensor, our initial tests showed that after storage 3 days at room temperature or 6 days in a fridge at 3-5 °C, the optical fiber biosensors functionalized with the pig IgG antibody still worked very well. When the storage time was longer than the indicated days, the activity of the fiber biosensor functionalized with pig IgG antibody decreased significantly.

5. Conclusion

In conclusion, an ultra-highly sensitive tapered SNSFC label-free biosensor for measurement of *S. aureus* is proposed and experimentally demonstrated. The tapered SNSFC sensor is composed of two parallel physical contact SNS fiber structures, where the two NCF sections are heated to be tapered to small diameter. The tapered SNSFC fiber sensor is then functionalized with pig IgG antibody which can specifically bind with *S. aureus*. Experimentally, for the tapered SNSFC sensor with a taper waist diameter of 10.4 μ m and functionalized with 200 μ g/ml pig IgG antibody, an average wavelength shift of 2.04 nm is obtained over a period of 30 minutes for the measurement of *S. aureus* with concentration of 7×10¹ CFU/ml. The tapered SNSFC has a large taper waist dimension (two 10.4 μ m diameter fibers), and hence has a good stability, which has been demonstrated by achieving a maximum wavelength variation of ±0.03 nm in the PBS for 40 minutes. The LOD of the biosensor is calculated as low as 3.1 CFU/ml, showing an ultrahigh sensitivity. The developed tapered SNSFC sensor structure can be used for various applications such as earlier diseases/medical diagnostics and food safety inspection once the fiber sensor is properly functionalized.

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